Cholesterotic Fibrous Histiocytoma in a Patient with Metabolic Syndrome

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Abstract
Among the many variants of dermatofibroma, dermatofibroma with cholesterol cleft (cholesterotic fibrous histiocytoma) is extremely rare. Here, we describe the case of a 50-year-old male patient with a cholesterotic fibrous histiocytoma on his left lower leg. He presented with a hyperkeratotic nodule 6 mm in diameter with a brown surface on the extensor surface of his left lower leg. The lesion had developed over the course of a few years without any tendency to heal. A skin biopsy performed on the tumor showed histopathological findings compatible with those of dermatofibroma. Interestingly, the lesion included many cholesterol clefts, as well as foamy histiocytes and multinucleated giant cells around them. He had had metabolic syndrome for years. To the best of our knowledge, this is the first report of a cholesterotic fibrous histiocytoma in a patient with metabolic syndrome. We conclude that the altered microenvironment caused by metabolic syndrome, as well as hyperlipoproteinemia itself, may play a role in the pathogenesis of this rare case.
Deguchi and Aiba: Cholesterotic Fibrous Histiocytoma in a Patient with Metabolic Syndrome

Introduction

Dermatofibroma has many variants, including cellular benign fibrous histiocytoma; angiomatoid-type, lipidized-variant, and granular cell dermatofibroma; epithelioid cell histiocytoma; atrophic dermatofibroma; and palisading fibrous histiocytoma. Among these variants, dermatofibroma with cholesterol cleft (cholesterotic fibrous histiocytoma) is one of the rarest forms. Histological findings of cholesterotic fibrous histiocytoma include cholesterol clefts within the center of the lesion, as well as variable degrees of invasion by foamy macrophages and giant cells around them. Furthermore, we could find only 3 definitive reports of cholesterotic fibrous histiocytoma associated with hyperlipoproteinemia. The first report was published by Hunt et al. [1] in 1990, followed by the reports by Yu et al. [2] and Takahashi et al. [3].

Recently, the concept of metabolic syndrome has been established. It was originally derived from the proposition by Reaven [4] named “syndrome X,” where atherogenic risk factors combine with underlying insulin resistance. Others have developed this concept into metabolic syndrome, defined by a cluster of pathological conditions including abdominal obesity, dyslipidemia, hyperglycemia, and hypertension.

Here, we describe the first case of cholesterotic fibrous histiocytoma associated with metabolic syndrome.

Case Report

A 50-year-old male presented with a hyperkeratotic nodule 6 mm in diameter with a brown surface on the extensor surface of his left lower leg (Fig. 1). The lesion had developed over the course of a few years without any tendency to heal. He had been suffering from assteatotic dermatitis on his four extremities, and admitted that he had scratched his lower legs frequently before the tumor developed. He was treated for hyperlipidemia and hyperglycemia, in addition to control of abdominal obesity. The laboratory examinations from half a year before demonstrated serum total cholesterol levels ranging from 344 to 479 mg/dL (reference interval: 125–220 mg/dL), a low-density lipoprotein cholesterol level between 236 and 311 mg/dL (reference interval: 70–140 mg/dL), a triglyceride level of 1,650 mg/dL at the maximum (reference interval: 45–150 mg/dL), and hemoglobin A1c ranging from 6.1 to 8.7% (reference interval: 4.3–5.8%).

A skin biopsy performed on the tumor showed acanthosis and basal hypermelanosis of the epidermis. The poorly demarcated tumor mass was located in the dermis, separated by a clear zone from the epidermis, extending into the deep reticular area (Fig. 2a). The tumor was composed of an admixture of fibroblast-like spindle cells, rounded histiocytes, and capillaries. Spindle cells were arranged irregularly in the collagen strands. There were neither atypical cells nor bizarre cells in the lesion. Within the center of the lesion there were abundant cleft spaces compatible with cholesterol clefts (Fig. 2b), surrounded by foam cells and giant cells. Immunohistochemistry of the tumor revealed CD68-positive cells as well as numerous factor XIIIa-positive cells, in contrast to a negativity for CD34. Based on these findings, we diagnosed this lesion as cholesterotic fibrous histiocytoma.

Later, we performed total excision of this tumor, and the patient has remained free from recurrence for more than 5 years.
Discussion

Our patient presented with dermatofibroma with many cholesterol clefts in the tumor mass, that is, cholesterotic fibrous histiocytoma. This case can be discriminated from tuberous xanthomas in that it had dermal cellular infiltrations by heterogeneous components including proliferative populations. They were composed of fibroblastic cells interspersed with collagen bundles and rounded histiocytic cells, in addition to foam cells and giant cell. Abundant factor XIIIa-positive cells in the tumor also supported the diagnosis of dermatofibroma in our case. Another differential diagnosis in this case includes lipidized fibrous histiocytoma. Although lipidized fibrous histiocytomas sometimes have cholesterol clefts, it is only a minor and infrequent finding [5] in their granulomatous lesions, whereas it is a predominant characteristic in our case; cholesterotic fibrous histiocytoma is defined as cholesterol deposition within the center of the lesion [2]. In addition, there was no hyalinized wiry collagen in the stroma of the specimen of this patient, which is known to be pathognomonic of lipidized fibrous histiocytoma [5].

The abdominal obesity, hyperglycemia, and hyperlipidemia (including hypertriglyceridemia) in our patient fulfilled the international criteria for metabolic syndrome [6, 7] as well as the Japanese standard [8]. To the best of our knowledge, this is the first case of cholesterotic fibrous histiocytoma in metabolic syndrome. The exact pathogenesis of cholesterol clefts remains to be elucidated. Yu et al. [2] proposed a hypothesis concerning the formation of cholesterol crystals in the lesion dermis; a possible local trauma may cause vascular leakage of lipoproteins followed by phagocytosis by dermal macrophages, resulting in such deposition. Actually, Takahashi et al. [3] demonstrated by observation with an electron microscope that in cholesterotic fibrous histiocytoma lesions, a cholesterol substance is incorporated in the histiocytes to form cholesterol crystals in the cytoplasm and lipid droplets in the lysosomes. Such a hypothesis seems to be most conceivable, since we can understand such a mechanism also by referring to xanthomas. As our patient had been suffering from itching of his lower legs due to asteatotic dermatitis, the scratching may have been responsible for minor injuries to the skin, leading to leakage of extremely high levels of serum cholesterol from the capillaries, as in reported cases of xanthoma [9, 10]. The accumulated cholesterol, especially low-density lipoprotein, may be altered by the microenvironmental oxidative stress, making it susceptible to phagocytosis [11, 12]. This speculation may be consistent with the common concept on the pathogenesis of atherosclerosis in the context of chronic inflammation including macrophage infiltration and oxidative stress [11–13]. Furthermore, the decreased level of plasma adiponectin found in metabolic syndrome patients [14] is well known to promote the uptake of oxidized low-density lipoprotein by macrophages, which may facilitate the formation of foam cells in the lesion. These microenvironmental conditions may combine to generate this rare variant of dermatofibroma. It is possible that with the increase in the number of cases with metabolic syndrome and hyperlipidemia in the general population, the number of patients with cholesterotic fibrous histiocytoma will increase as well. An analysis of case reports such as ours may enable us to predict the prognosis of metabolic syndrome and to elucidate the exact pathomechanism of cholesterol clefts themselves.
Statement of Ethics

We confirm that the patient provided written informed consent to use his photographs for publication.

Disclosure Statement

The authors have nothing to disclose.

References

Fig. 1. Clinical appearance of dermatofibroma in the present patient.

Fig. 2. a Histological findings from a biopsy specimen of the tumor. Scale bar, 200 μm. b Intratumoral layers of cholesterol clefts. Scale bar, 200 μm.